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Application for U.S. Letters Patent Entitled

PROTEIN COMPLEXES HAVING FACTOR VIII:C ACTIVITY AND PRODUCTION THEREOF

which is a continuation of U.S. patent application serial no. 10/256,849 filed September 26, 2002, which is a continuation of 09/748,062 filed December 22, 2000, which is a continuation of U.S. patent application serial no. 08/441,943 filed May 16, 995 (USP 6,228,620), which is a divisional of U.S. patent application serial no. 08/161,770 filed December 3, 1993 (USP 5,595,886), which is a continuation of U.S. patent application serial no. 07/652,099 filed February 7, 1991, which is a continuation-in-part of U.S. patent application serial no. 07/051,916 filed May 19, 1987, which is a continuation-in-part of U.S. patent application serial no. 06/822,989 filed January 27, 1986.

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Attorney Docket No. PP00048.252 (2300-0048.05)

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PROTEIN COMPLEXES HAVING FACTOR VIII:C <u>ACTIVITY AND PRODUCTION THEREOF</u>

Cross-reference to Related Application

This application is a continuation-in-part of copending U.S. Patent Application Serial No. 051,916, filed 19 May 1987, which is a continuation-in-part of U.S. Patent Application Serial No. 822,989, filed 27 January 1986, now abandoned, both of which are incorporated herein by reference in full.

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Description

Technical Field

This invention relates to protein complexes having Factor VIII:C activity, and to methods for producing said complexes by expression of suitable polynucleotide constructs. The protein complexes are useful in the treatment of classical (Type A) hemophilia.

Background of the Invention

Hemophilia A is an X-chromosome-linked inherited disease which afflicts 1-2 males per 10,000. The disease is caused by an absence or deficiency of Factor VIII:C. Factor VIII:C is a very large glycoprotein (native M_r 330 K - 360 K), which is present in plasma at extremely low concentrations. It

is a necessary element in the proteolytic cascade which converts soluble fibrinogen to insoluble fibrin, forming a clot to prevent blood loss from traumatized tissue. In the bloodstream, it is found in noncovalent association with Factor VIII:R ("von Willebrand factor"), which acts as a stabilizing carrier protein. Factor VIII:C is very susceptible to cleavage by thrombin, plasmin, protease C, and other serine proteases. It is generally isolated from plasma or plasma products as a series of related polypeptides ranging from M_r 160 K-40 K with predominant species of M_r 92 K and M_r 80 K-77 K. This complex pattern has made the analysis of the structure of active Factor VIII:C very difficult.

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Factor VIII:C and the related polypeptides have been described by F. Rotblat et al, Biochemistry (1985) 24:4294-4300; G.A. Vehar et al, Nature (1984) 312:337-342; J.J. Toole et al, Nature (1984) 312:342-347; and M.A. Truett et al, DNA (1985) 4:333-349. E. Orr et al, Molecular Genetics of Clotting Factors, p. 54, s321, reported a "spacer" function for the heavily glycosylated region of Factor VIII:C. The sequence has been reported by J.J. Toole et al, supra; W.I. Wood et al, <u>Nature</u> (1984) <u>312</u>:330-336; and M.A. Truett et al, supra. full-length protein contains three repeats of one sequence (I), and two repeats of a second sequence (III). A third, heavily glycosylated sequence (II) is present between the second and third I repeats, and is apparently cleaved proteolytically to form the M_r 92 K and M_r 80 K polypeptides. The first two I repeats form the A domain, while the third I repeat and the two III repeats form the C domain. The II sequence forms the B domain. Thus, the full-length protein has the structure I,-I2-II-I3-III1-III2 (A-B-C), while the Mr 92 K and Mr 80 K polypeptides (A and C) have the structures I_1-I_2 and $I_3-III_1-III_2$, respectively. C. Fulcher et al, <u>J Clin Invest</u> (1985) <u>76</u>:117-124, suggested that based on antibody-epitope data with Factor

VIII:C, both the M_r 92 K and the M_r 80 K polypeptides are necessary for Factor VIII:C function.

Factor VIII:C has historically been isolated from blood in a concentrated form for therapeutic treatment of hemophilia. However, concerns regarding transmission of HIV and other blood-borne diseases have stimulated activity to provide alternative supplies of Factor VIII:C. It is of substantial interest to be able to supply compositions having Factor VIII:C activity without concerns as to the transmission viral diseases associated with the native Factor VIII:C.

Although full-length recombinant human Factor VIII:C has been produced, it is difficult to purify and characterize, and it is unstable due to proteolysis. Efficient recombinant production of the full-length molecule for clinical use is doubtful at this time.

R.L. Burke et al, <u>J Biol Chem</u> (1986) <u>261</u>:12574-78 disclosed the expression of an active Factor VIII:C complex from cells simultaneously transfected with polynucleotides encoding M_r 92 K and M_r 80 K polypeptides. The obtained protein demonstrated activity equal to that of cloned full-length Factor VIII:C expressed under similar conditions. O. Nordfang et al, <u>J Biol Chem</u> (1988) <u>263</u>:1115-18 disclosed the *in vitro* assembly of active Factor VIII:C complexes from separate preparations of M_r 92 K protein and M_r 80 K protein (FVIII-HC and -LC, respectively). Successful assembly required divalent metal ions (especially Mn⁺⁺ and Ca⁺⁺) and thiols, but only a small amount of FVIII-HC could be complexed into active FVIII:C.

30 <u>Disclosure of the Invention</u>

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We have now invented an improved method for expressing recombinant protein complexes with high stability and Factor VIII:C activity. The $M_{\rm r}$ 92 K polypeptide (FVIII-HC) and the $M_{\rm r}$ 80 K polypeptide (FVIII-LC) are expressed as two separ-

ate polypeptides, under the control of separate promoters, within the same host cell. Each polypeptide is preferably expressed using a signal sequence which directs export to the extracellular space with cleavage of the signal sequence. FVIII-HC is preferably expressed as a fusion protein having a 5 C-terminal extension. The extension comprises a polypeptide sequence homologous to the B domain N-terminal sequence (which may allow cleavage by thrombin), a polypeptide spacer of 3 to 100 amino acids, and a sequence homologous to the C-terminal B domain sequence. The C-terminal extension of FVIII-HC results 10 in a higher yield of active polypeptide upon expression in eukaryotic host cells. FVIII-LC is preferably expressed as an LC polypeptide using a signal peptide. The FVIII-LC polypeptide is processed and secreted efficiently with the correct Nterminal amino acid residue, and correct glycosylation. 15 transfection with polynucleotides encoding FVIII-HC and FVIII-LC in a suitable host cell provides recombinant protein complexes having Factor VIII:C activity in high yield.

20 Modes of Carrying Out The Invention

A. <u>Definitions</u>

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The term "polynucleotide" as used herein refers to a sequence of DNA or RNA, which may be single or double-stranded (ss or ds), or a DNA-RNA heteroduplex. In most cases herein, polynucleotide will refer to dsDNA.

The term "signal peptide" as used herein refers to a peptide sequence which is recognized and acted upon by signal peptidase during expression of the polypeptide. Signal peptides encode peptide sites for signal peptidase cleavage, and cause the attached polypeptide to be transported into the secretion pathway leading to the extracellular medium.

The term "A domain" refers to that portion of human Factor VIII:C which constitutes the $M_{\rm r}$ 92 K protein subunit. The A domain contains from about 740 to about 760 amino acids,

and is found at the N-terminus of the native human Factor VIII:C. The A domain polypeptide will extend from amino acid 10, usually amino acid 1, to at least about amino acid 620, usually at least about amino acid 675, more usually at least about amino acid 740. The polypeptide will include at least about 85% of the A domain (Wood et al, supra), more usually at least about 90% and may optionally include a portion of the N-terminus of the B domain, typically not exceeding about amino acid 1405. Of particular interest is an N-terminal chain having the entire sequence to the thrombolytic cleavage site at Arg -Ser...

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The term "B domain" refers to that portion of native human Factor VIII:C which is generally removed by intracellular cleavage, and which is heavily glycosylated when expressed in mammalian cells such as COS7 and CHO. The B domain contains an N-terminal sequence, which allows cleavage of the A domain from the B domain by thrombin. The B domain also has a C-terminal processing site which allows cleavage of the C domain from the A-B precursor by an enzyme located in the Golgi apparatus of the mammalian cell. The sequences of the N-terminal and C-terminal sequences are set forth in the Examples below. The complexes of the invention which lack "a substantial portion of the B domain" lack all of the B domain, or essentially all of the B domain except for the N-terminal and C-terminal sequences.

The term "C domain" refers to that portion of native human Factor VIII:C which constitutes the C-terminus of the full length protein, and is cleaved intracellularly to form the Factor VIII:C light chain. The light chain will have an amino acid sequence substantially the same as the amino acid sequence of the C-terminus of a Factor VIII:C polypeptide, usually at least about 80%, more usually at least about 90% of the Factor VIII:C M_r 80 K chain, particularly beginning with amino acid 1570, usually amino acid 1600, particularly amino acid 1625,

more particularly amino acid 1640, preferably at about amino acid 1649, ±10 amino acids, more particularly ±1 amino acid, and continuing to at least about amino acid 2300, usually 2310, ±10 amino acids, preferably 2325, ±5 amino acids, more preferably to the terminal amino acid (2332). Usually, the light chain will have at least about 85%, more usually at least 95%, of the C1-C2 domains, desirably the A3-C1-C2 domains.

The term "co-expressing" as used herein refers to simultaneous expression of an A domain polypeptide and a C domain polypeptide within the same host cell. The polynucleotide sequences encoding the A and C domains may be on the same or on different expression cassettes or plasmids. Co-expression of the A and C domains permits proper folding to occur, which in turn provides an A-C complex having higher activity and efficiency of secretion.

The term "cell growth medium" as used herein refers to any medium suitable for culturing host cells, and includes media suitable for obtaining expression of recombinant products whether actual cell "growth" occurs or not. Cell growth media generally include nutrients and a metabolizable energy source in an aqueous solution. If desired, cell growth media may also include a compound which induces expression of the recombinant polypeptides of the invention. Selection of such an inducing compound depends upon the promoter selected to control expression. Other typical additives include selection compounds (i.e., drugs or other chemicals added to the media to insure that only transformed host cells survive in the medium) and serum, such as fetal bovine serum (FBS). "Serum-free medium". is a solution which has been supplemented to such an extent that the necessary trace factors present in serum need not be added in the form of serum. There are many suitable cell growth media available from commercial sources.

The term "polypeptide spacer" refers to a polypeptide sequence of about 3 to about 100 amino acids, which is gener-

ally not homologous to the human Factor VIII: C B domain, and which carries fewer than 5 potential sites of N-linked glycosylation. Preferably, there will be 2 or fewer such sites. is presently believed that the large size and high degree of glycosylation of the B domain prevents efficient expression of the $M_{\rm r}$ 92 K polypeptide. However, a low (but useful) yield of the Mr 92 K polypeptide is obtained when the B domain is completely removed. It is also presently believed that the A domain may not be folded correctly on a consistent basis in the absence of the B domain, so that only a small percentage of the A domain is correctly folded and expressed. The polypeptide spacer of the invention provides a C-terminal extension to the A domain, and apparently stabilizes the polypeptide and improves secretion in active form. Thus, it may be that use of a polypeptide which is glycosylated lightly (or not at all) prevents the A domain-spacer construct from encountering the same size problems obstructing expression of full-length Factor The presently preferred spacer is derived from a human Ig heavy chain hinge, particularly from human IgA1. spacer provides a flexible extension, without adding an immunogenic epitope (when administered in humans).

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The term "homology" as used herein means identity or substantial similarity between two polynucleotides or two polypeptides. Homology is determined on the basis of the nucleotide or amino acid sequence of the polynucleotide or polypeptide. In general terms, usually not more than 10, more usually not more than 5 number%, preferably not more than about 1 number% of the amino acids in the chains will differ from the amino acids naturally present in the Factor VIII:C A and C domains. Particularly, not more than about 5%, more usually not more than about 1% will be nonconservative substitutions. Conservative substitutions include:

Gly + Ala; Val + Ile + Leu;
Asp + Glu; Lys + Arg;

Asn + Gln; and

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Phe + Trp + Tyr.

Nonconservative changes are generally substitutions of one of the above amino acids with an amino acid from a different group (e.g., substituting Asn for Glu), or substituting Cys, Met, His, or Pro for any of the above amino acids.

The term "sufficient amount" of protein complex of the invention refers to that amount of protein which is capable of effecting therapeutic treatment of a subject having a disorder treatable with native human Factor VIII:C. In general, the protein complex of the invention is essentially as active as native human Factor VIII:C, and may be administered in similar amounts. The specific activity of the protein complex of the invention may be determined by means known in the art, as described below (e.g., by using the commercially available COATEST assay).

The term "effective concentration" refers to a concentration of expression cassette which is capable of transforming a host cell under appropriate transformation conditions.

B. <u>General Method</u>

DNA constructs are generally employed for expression of the polypeptides of the invention. Each of the polynucleotide constructs will have, in the 5'-3'-direction of transcription, a transcriptional initiation and translational initiation region, a structural gene coding region comprising a sequence coding for the signal peptide sequence, and a sequence coding for the Factor VIII:C heavy or light chains, followed by translational and transcriptional termination sequences. The selection of specific elements such as these is within the skill of the art.

The initiation region may comprise a number of different sequences related to the initiation of transcription and translation. These sequences include enhancer sequences, RNA polymerase binding site, RNA capping site, ribosomal binding and translational initiation sites, and the like. The transcriptional initiation region may be the natural region associated with Factor VIII:C, or may be an alternative sequence to provide for higher transcriptional efficiency. The sequences may be obtained from mammalian viruses or the genes of the host cell or genes from a different mammalian host which are active in the host cell. Numerous transcriptional initiation regions have been isolated and demonstrated to be operative in mammalian host cells. These regions include the SV40 early promoter and late promoter region, the adenovirus major late promoter region, actin promoter region, the cytomegalovirus M_r 72 K immediate early protein promoter region, the metallothionein promoter, and the like.

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The termination region may include 3'-untranslated sequences, a polyadenylation signal sequence, and the like. The termination region may be obtained from the 3' non-translated sequence of the Factor VIII:C natural cDNA, or may be from the same structural gene or different structural gene from which the 5'-initiation region was obtained. The 3'-region is not as essential to the level of transcription as the initiation region, so that its choice is more of a matter of convenience than specific selection.

The structural genes typically include a leader sequence coding for the signal peptide which directs the polypeptide into the lumen of the endoplasmic reticulum for processing and maturation. Optionally included are additional sequences encoding propeptides which are processed post-translationally by endopeptidases, where the endopeptidases cleave a peptide bond, removing the propeptide to generate the mature polypeptide. The signal peptide may be the naturally occurring one, particularly for the N-terminal peptide, or may be any signal peptide which provides for the processing and maturation of the polypeptides.

Various signal peptides have been reported in the literature and include such sequences as that of tissue plasminogen activator, immunoglobulin heavy and light chains, viral membrane glycoproteins such as Herpes Simplex virus glycoproteins gB and gD, α_1 -antitrypsin, and the like. The α_1 -antitrypsin signal peptide is presently preferred for secretion of the FVIII-LC polypeptide.

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The DNA sequences encoding the mature protein and signal peptide must be joined so as to be in reading frame. Where convenient restriction sites are available, the cohesive or blunt ends may be properly joined. However, for the most part, adapters will be employed where portions of the coding sequence will be recreated in the synthetic adaptor so that the truncated structural gene and/or truncated signal sequence will be linked through the adaptor, so as to be in proper reading The signal sequence and structural gene may be partially restriction mapped, so as to identify restriction sites, particularly unique restriction sites, which may be employed to link the two sequences together in proper reading frame by means of an appropriate adaptor. Alternatively unique restriction sites may be inserted at the junction of the signal sequence and mature polypeptide coding sequence by in vitro mutagenesis.

The translational start and stop signals will normally be part of the structural gene, providing for the initiation codon at the beginning of translation and one or more stop codons for the termination of translation. The initiation codons will be the first codons of the signal sequences. The stop codons may be added as appropriate as part of the termination region or be added to the coding region to provide for convenient 3'-terminus for linkage to the transcriptional termination region to provide for a complete termination region.

The various regions of the expression cassette, (the transcriptional and translational initiation region nucleic

acid sequence, structural gene nucleic acid sequence encoding one of the polypeptides and under the transcriptional and translational control of the initiation region, and a transcriptional and translational termination region, controlling the processing of the mRNA and the translational termination) which identify the particular nucleotide sequences may be joined using conventional methods. Usually, the sequences obtained will contain, or be modified to contain restriction sites, which may then be annealed where complementary overhangs or cohesive ends are present. Modification frequently will be in noncoding regions by the introduction of linkers to provide for the desired cohesive ends. The ends will usually be ligated prior to introduction into the host cell, although the host cell may be allowed to provide the necessary ligation.

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The expression cassettes may be joined to a wide variety of other sequences for particular purposes. Where amplification of the amount of secreted glycoprotein is desired, the expression cassettes for FVIII:C may be joined in tandem to a gene for which spontaneous increases in gene copy number can be selected by an appropriate treatment. Such genes include the human metallothionein gene, and the mouse dihydrofolate reductase gene. These genes are placed in cassettes having their own transcriptional and translational regulatory sequences. By selecting cell clones resistant to increasing concentrations of heavy metal ions (e.g., cadmium) or methotrexate, the gene of interest (the expression cassette) may be co-amplified in the host cell.

The subject expression cassettes may be part of a vector comprising a replication system functional in the host cell, which replication system may provide for stable episomal maintenance or integration of the expression cassette into the host genome. The vector will also comprise a marker for selection, for selecting mammalian host cells containing the DNA

construct and the vector from those host cells which lack the DNA construct and vector.

A wide variety of replication systems are available, typically derived from viruses that infect mammalian host cells. Illustrative replication systems include the replication systems from Simian virus 40, adenovirus, bovine papilloma virus, polyoma virus, Epstein Barr virus, and the like.

Selection markers enabling propagation of the vector in prokaryotic host cells may include resistance to a biocide, particularly an antibiotic, or complementation of auxotrophy to provide a prototrophic host. Particular genes of interest as markers include kanamycin resistance gene (NPTII), chloramphenicol resistance gene (CAT), penicillinase (β -lactamase), or the like.

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one or more restriction sites which allow for the insertion of the expression cassette, stepwise or as a completed entity, into the vector. Frequently, the vector will also include a bacterial replication and selection system, which allows for cloning after each of the manipulative steps. In this way, relatively large amounts of the construction at each of the stages may be prepared, isolated, purified, tested to verify that the proper joining has occurred, and then used for the next stage.

Various mammalian host cells may be employed in which the regulatory sequences and replication system are functional. Such cells include COS7 cells, Chinese hamster ovary (CHO) cells, mouse kidney cells, hamster kidney cells, HeLa cells, HepG2 cells, or the like.

The expression cassettes of the desired polypeptides may be linked together in one nucleic acid chain or may be provided in separate nucleic acid molecules. The expression cassettes may be parts of different vectors or of the same vector. This is primarily a matter of convenience, although in some

situations with particular vectors, one or the other manner of construction may be preferable.

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The expression vector may be a replication-deficient retrovirus. S.-F. Yu et al, Proc Nat Acad Sci USA (1986) 83: 3194-98 disclosed the construction of self-inactivating ("SIN") retroviral gene transfer vectors. SIN vectors are created by deleting the promoter and enhancer sequences from the U3 region of the 3' LTR. A functional U3 region in the 5' LTR permits expression of the recombinant viral genome in appropriate packaging cell lines. However, upon expression of its genomic RNA and reverse transcription into cDNA, the U3 region of the 5' LTR of the original provirus is deleted, and is replaced with the U3 region of the 3' LTR. Thus, when the SIN vector integrates, the non-functional 3' LTR U3 region replaces the functional 5' LTR U3 region, and renders the virus incapable of expressing the full-length genomic transcript.

The expression cassettes are introduced into the host cell by conventional methods. Conveniently, calcium phosphate-precipitated DNA or DNA in the presence of DEAE-dextran may be employed for transformation. A synthetic lipid particularly useful for polynucleotide transfection is N-[1-(2,3-dioleyl-oxy)propyl]-N,N,N-trimethylammonium chloride, which is commercially available under the name Lipofectin® (available from BRL, Gaithersburg, MD), and is described by P.L. Felgner et al, Proc Nat Acad Sci USA (1987) 84:7413. Where viruses are involved, transfection or transduction may be employed. The particular manner in which the host cell is transformed is not critical to this invention, depending substantially upon whether the expression cassettes are joined to a replication system and the nature of the replication system and associated genes.

The transformed/transfected cells are then grown in an appropriate nutrient medium. It is presently preferred to employ CHO cells cultured at 10 to 32°C, more preferably about

27°C, for less than about 30 hours, more preferably less than 10, most preferably less than about 4 hours. The product is obtained as a complex of the two FVIII:C chains, so that the media or cell lysate may be isolated and the Factor VIII:C active complex extracted and purified. Various means are available for extraction and purification, such as affinity chromatography, ion exchange chromatography, hydrophobic chromatography, electrophoresis, solvent-solvent extraction, selective precipitation, and the like. The particular manner in which the product is isolated is not critical to this invention, and is selected to minimize denaturation or inactivation and maximize the isolation of a high-purity active product.

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Compositions are provided where the composition in the COATEST assay will have at least 0.02 U/mL of activity, usually at least about 0.2, more usually at least about 0.5 U/mL of activity. The subject product can be purified by affinity chromatography using antibodies, particularly monoclonal antibodies directed against the FVIII-LC, electrophoresis, extraction, HPLC, etc.

The subject method provides for production of a complex of the heavy and light chains which has Factor VIII:C activity. Production is evidenced by conditioned media as described in the experimental section, which will have at least about 50, usually at least about 70 mU/mL, more usually at 25 least about 200 mU/mL of Factor VIII:C activity in the COATEST assay.

The complexes having Factor VIII:C activity produced according to the invention have a variety of uses as immunogens for the production of antibodies, for isolation of von Willebrand factor by affinity chromatography, in diagnostic assays for Factor VIII:C and for treatment of hemophiliacs and other hosts having blood clotting disorders. The subject protein complexes may be administered in a physiologically acceptable carrier, such as water, saline, phosphate buffered saline, and

citrate buffered saline, at concentrations in the range of about 10-200 U/mL. See U.S. Patent Nos. 3,631,018; 3,652,530, and 4,069,216 for methods of administration and amounts. Other conventional additives may also be included.

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C. <u>Examples</u>

The examples presented below are provided as a further guide to the practitioner of ordinary skill in the art, and are not to be construed as limiting the invention in any way.

Example 1

(Preparation of Expression Plasmids)

(A) pSV7d:

The expression cassettes were prepared using the mammalian cell expression vector pSV7d (2423 bp).

The plasmid pSV7d (see Truett et al, supra) was constructed as follows: The 400 bp BamHI/HindIII fragment containing the SV40 origin of replication and early promoter was excised from pSVgtI (obtained from Paul Berg, Stanford University, California) and purified. The 240 bp SV40 BclI/BamHI fragment containing the SV40 polyA addition site was excised from pSV2/DHFR (Subramani et al, Mol Cell Biol (1981) 1:854-864) and purified. The fragments were fused through the following linker:

Stop Codons

1 2 3

5'-AGCTAGATCTCCCGGGTC<u>TAGATAAGTAA</u>T-3'
TCTAGAGGGCCCAGATCTATTCATTACTAG

HindIII BglII Smal Xbal

BclI overhang.

This linker contains five restriction sites, as well as stop codons in all three reading frames. The resulting 670 bp fragment containing the SV40 origin of replication, the SV40 early promoter, the polylinker with stop codons and the SV40 polyadenylation site was cloned into the BamHI site of pML, a pBR322 derivative having about 1.5 Kb deleted (Lusky and Botchan, Cell (1984) 36:391), to yield pSV6. The EcoRI and EcoRV sites in the pML sequences of pSV6 were eliminated by digestion with EcoRI and EcoRV, treated with Bal31 nuclease to remove about 200 bp on each end, and finally religated to yield The Bal31 resection also eliminated one BamHI restriction site flanking the SV40 region, approximately 200 bp away from the EcoRV site. To eliminate the second BamHI site flanking the SV40 region, pSV7a was digested with NruI, which cuts in the pML sequence upstream from the origin of replication. This was recircularized by blunt end ligation to yield pSV7b.

pSV7c and pSV7d represent successive polylinker replacements. First, pSV7b was digested with StuI and XbaI. Then, the following linker was ligated into the vector to yield pSV7c:

Thereafter, pSV7c was digested with BglII and XbaI, and then ligated with the following linker to yield pSV7d:

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(B) pSVF8-92:

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pSVF8-92 is an expression plasmid for the Mr 92 K Starting from the BamHI site in the polylinker FVIII-HC chain. pSV7d, pSVF8-92 consists of a 49 bp synthetic linker-adaptor molecule from BamHI to SacI encoding nucleotides -30 to +14 of the Factor VIII: C cDNA, (numbering from the first A of the translational start site; the sequence is shown below in (D) a way and a 2267 bp SacI to HindIII fragment from the Factor VIII:C DNA contained in pSVF8-200 described below (up to nucleotide +2281), and pSV7d from HindIII to BamHI.

(C) pSVF8-80:

pSVF8-80 is an expression plasmid for the M_r 80 K FVIII-LC chain. Starting from the SalI site in the polylinker pSV7d, pSVF8-80 consists of a 201 bp fragment of a tissue plasminogen activator cDNA from nucleotides -98 to +103 (relative to the start codon) terminating at a BglII site (tPA sequences given in S.J.F. Degan et al, J Biol Chem (1986) 261:6972-6985), a 29 bp synthetic BglII to BclI linker-adaptor encoding nucleotides +5002 to +5031 of Factor VIII:C ligated to a 2464 bp BclI fragment of factor VIII:C spanning from a BclI site created at nucleotide 5028 of the factor VIII:C cDNA through in vitro mutagenesis (Zoller and Smith, Meth Enzymol (1983) 100:468) (pF8GM7), to a BclI site in the 3' untranslated region, at nucleotide 7492, and a 400 bp fragment of tPA 3' untranslated sequence spanning from a BglII site to a synthetic PstI site generated from the cDNA cloning, followed by the polylinker from the vector M13mp9 (Vieira and Messing, Gene (1982) 19:259) and then pSV7d.

(D) pSVF8-200

The vector pSVF8-200 is an expression plasmid for the full-length Factor VIII: C cDNA. The plasmid pSVF8-200 (described in Truett et al), which contains the entire Factor

VIII:C cDNA coding and 3' untranslated sequences, with the 5' untranslated sequences the same as described above for psvF8-92, was prepared as follows.

Plasmid pSV7d was digested with BamHI to cut in the polylinker region downstream of the SV40 early promoter. The following 49 bp BamHI-SacI linker adaptor, which codes for the last 30 bp of the 5' untranslated region and the first 15 bp of the human Factor VIII:C coding sequence, was chemically synthesized and ligated to pSV7d.

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GATCC TCTCC AGTTG AACAT TTGTA GCAAT AAGTC

BamHI G AGAGG TCAAC TTGTA AACAT CGTTA TTCAG

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Met Gln Ile Glu ATG CAA ATA GAG CT 3 TAC GTT TAT C<u>Sac</u>I 5

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This ligated plasmid was subsequently digested with SacI to remove excess linkers and with SalI to provide a SalI overhang.

Fragment 1, the 2.9 K SacI fragment from pF8-102 containing the 5' coding region of human Factor VIII:C, and Fragment 2, the 6.5 K SacI-SalI fragment from pF8-6.5 which contains the 3' coding region of the factor, and pSV7d modified vector containing the linker adaptor were ligated together (see Truett et al, supra). This ligation mix was then used to transform E. coli HB101, and colonies were selected by resistance to ampicillin.

Three hundred transformants were screened by colony filter hybridization using the BamHI-SacI 5' adaptor or the 2.9 K SacI fragment as probes. Those colonies positive with both probes were then analyzed by restriction mapping. Plasmid pSVF8-200, which contains the entire coding region for the human Factor VIII:C gene and a 5' untranslated region properly

fused in transcriptional orientation to the SV40 early promoter, was obtained.

(E) Transfection and Culture of COS7 Cells:

The plasmids described above were transfected into COS7 cells (Guzman, Cell (1981) 23:175) using the calcium phosphate coprecipitation method (van der Eb and Graham, Meth Enzymol (1980) 65:826-39) coupled with treatment with chloroquine diphosphate (Luthman and Magnusson, Nuc Acids Res (1983) 11:1295-1308) using 50 µg of plasmid DNA per 5×10⁵ cells for 14 hr. Cells may also be transfected by the DEAE-dextran method of Sompayrac and Danna, Proc Nat Acad Sci USA (1981) 78:7575-78.

The COS7 cells were cultured in Dulbecco's modified Eagle medium supplemented with 10% fetal calf serum, 100 U/mL penicillin, 100 μ g/mL streptomycin, 292 μ g/mL glutamine, and 110 μ g/mL sodium pyruvate. Samples were obtained from a 48-hour collection of serum-containing medium at 88 hours post transfection.

(F) Assays

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At specific intervals post transfection, medium was removed from the cells, and aliquots were stored at -70°C. Samples were tested for their ability to decrease the prolonged partial thromboplastin time of Factor VIII:C deficient plasma in a standard coagulation assay (Hardisty et al, Thromb et Diathesis Haemolog (1962) 72:215). The more specific COATEST assay (Rosen et al, Thromb and Haemostasis (1985) 54:818-823), which measures the generation of activated Factor X (Xa) as a linear function of the concentration of exogenously supplied Factor VIII:C, was used to verify the results of the coagulation assay. The concentration of immunologically reactive Factor VIII:C protein in the medium was determined by the application of a radioimmunoassay (RIA) developed to detect the M, 92

K polypeptide and by an enzyme-linked immunosorbant assay (ELISA) specific for the M_r 80 K polypeptide (Nordfang et al, Thromb Haemostasis (1985) 53:346).

As shown in Table 1, expression of the M_r 92 K polypeptide or of the M_r 80 K polypeptide alone produced no detectable activity even though high levels of each of the individual proteins were present in the conditioned media. When cells were cotransfected with pSVF8-92 and pSVF8-80 plasmids, the media contained about 20 mU/mL of coagulation activity. The same relative level of the coagulation activity was secreted by cells transfected with the plasmid pSVF8-200 encoding the complete Factor VIII:C protein.

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When conditioned media from the pSVF8-92 and the pSVF8-80 single transfectants were mixed together (using several different conditions as outlined in Table 1) no activity was measurable.

These results indicate that a complex of the amino and carboxyl terminal domains of Factor VIII:C retains intrinsic coagulation activity and that the interior domain is not essential for activity nor for the assembly of an active complex from separate chains.

TABLE 1: Assay of Recombinant Factor VIII:C Activity in Conditioned COS7 Cell Media

5	Plasmid		lation ''' Activity mU/mL	COATEST Activity mU/mL	HC-RIA Assay U/mL	LC-ELISA Assay U/mL	
10 :	pSVF8-92	95.7	<0.9	<0.1	0.15	<0.0002	er egizil (desire sense)
	pSVF8-80	97.2	<0.9	<0.1	<0.01	1.36	
15	pSVF8-92 & pSVF8-80a	56.1	22.5	20.4	0.05	1.13	
	pSVF8-200	47.7	70.0	43.2	. 0.12	0.28	
20	none	94.6	<0.9	<0.1	<0.0	<0.0002	
20	pSVF8-92J+ pSVF8-80b†	95.7	<0.9	<0.1	_	<u>-</u>	. •

plasmids were cotransfected into the same cells
plasmids were transfected into separate cells, and the
supernatants mixed 48 hours later

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In Table 1, Coagulation Time and Activity were obtained as follows: Aliquots of 75 μ L of media, conditioned by the growth of COS7 cells transfected with the indicated plasmids or mock transfected, were assayed for their ability to decrease the prolonged partial thromboplastin time of Factor VIII:C-deficient plasma in the one-stage assay. Briefly, 75 μ L of Platelin (General Diagnostics) was incubated for 3 min at 37°C, followed by the addition of 75 μ L of Factor VIII:C-deficient plasma plus 75 μ L of the test sample for an additional 5 min incubation at 37°C. A 75 μ L aliquot of prewarmed 0.025 M CaCl, was added, and the clotting time measured with a Becton-

A variety of mixing conditions were tested, including preincubation for various times up to 2 hr at 37°C, 20°C, or 4°C in the presence or absence of 10 mM CaCl₂. The value reported in this table is representative of the data obtained.

Dickinson fibrometer. Normal human plasma diluted in COS7 cell medium was used as a standard. One mU of activity is assumed to correspond to approximately 100 pg of Factor VIII:C protein (Fay et al, Proc Nat Acad Sci USA (1982) 79:7200).

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In Table 1, the COATEST assay (Kabi) was used to measure the generation of activated Factor X (Xa) as a linear function of the concentration of Factor VIII: C. The concentration of Factor Xa is measured by the proteolytic cleavage of the chromogen para-nitroaniline from a synthetic peptide substrate for Factor Xa. Normal human plasma diluted in 50 mM Tris-HCl, pH 7.3, 0.2% BSA was used as the standard.

For the RIA assay in Table 1, purified canine Factor VIII: C-inhibitory IgG was coated onto the wells of a 96-well polystyrene microtiter plate at a concentration of 3.5 μ g/mL in 0.1 M sodium carbonate buffer, pH 9.8, by overnight incubation The plates were washed 3 times with 0.1 M NaCl, 0.05% Tween® 20 followed by incubation with a mixture of test medium samples and iodinated FVIII:C M, 92 K protein, both diluted in 0.05 M imidazole, 0.1 M NaCl, 1% bovine serum albumin, 0.05% Tween® 20, pH 7.3. The FVIII:C M, 92 K protein was isolated from plasma and was greater than 50% homogeneous as estimated by SDS-PAGE and silver staining. After incubation for 16 hr at room temperature, the plates were washed, and the amount of 125I in the individual wells was measured in a gamma counter. An intermediate purified commercial Factor VIII:C preparation (Factor VIII, NORDISK) with a specific activity of 0.5 unit of coagulation activity per mg was used as the standard. standard was calibrated against the World Health Organization Third International Factor VIII:C standard. We defined our intermediate purified standard to contain a M, 92 K RIA activity/Factor VIII:C coagulation activity ratio of 1.

For the ELISA assay in Table 1, purified human Factor VIII:C-inhibitory IgG was coated onto the wells of a 96-well PVC microtiter plate at a concentration 4.5 μ g/mL in 0.1 M

sodium carbonate, pH 9.8, by overnight incubation at 37°C. The wells were washed as above and peroxidase-conjugated F(ab')₂ fragments of the human inhibitory IgG diluted in 0.1 M imidazole, 0.15 M NaCl, 1% BSA, 0.05% Tween® 20, pH 7.3, were added for a final incubation of 16 hr at room temperature. The color was developed with o-phenylenediamine solution. Normal human serum was used as a standard.

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To verify that the observed coagulation activity was due to Factor VIII:C, the sensitivity of the coagulation to inhibition by antibody specific for Factor VIII:C was determined. Prior to assay, aliquots of conditioned media were preincubated for 2 hr at 37°C in the presence of dilutions of normal human serum or of serum from a hemophiliac who had developed a high titer of inhibitory antibodies to Factor VIII:C. As shown in Table 2, the activity of the complete molecule, as well as that of the Mr 92 K-80 K complex was reduced specifically by the inhibitory serum. The same results were obtained using three different inhibitory monoclonal antibodies which bind to the Mr 80 K species. Inhibition of Factor VIII:C activity using inhibitory serum was studied as follows: of the indicated COS7 cell conditioned medium were incubated with 20 µL of a 100-fold dilution of human Factor VIII:C inhibitory serum (Bethesda titer 1500 units) or a similar dilution of pooled normal human serum, or buffer alone (50 mM imidazole, 0.1 M NaCl, 100 μ g/mL BSA pH 7.3) for 2 hr at 37°C. samples were then assayed for residual coagulation activity as outlined above.

Coagulation Inhibition Assay

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Plasmid	serum	Coagulation Time (secs)
pSVF8-80 + pSVF8-92	Normal	. 51.9
,- 	Immune	74.5
nga ngabantar i mga salah na daga mga	Buffer	54.4
pSVF8-200	Normal	46.4
	Immune	69.4
	Buffer	46.8

The inhibition experiment was repeated using monoclonal antibodies, as follows: 100 μ L of conditioned medium were incubated for 2 hr at 37°C with either 10 μ L of a 1 μ g/ μ L solution of anti-Factor VIII: C monoclonal antibody from Hybritech (Bethesda titer 14,000 units) or buffer, and then assayed as above. The results are shown in Table 3.

Coagulation Inhibition Assay

Plasmid	serum	Coagulation Time (secs)
pSVF8-92 + pSVF8-80	Immune	72.9
-	Buffer	48.0
pSVF8-200	Immune	60.9
	Buffer	44.9

To demonstrate more clearly the existence of a two chain complex, the active species was partially purified from the COS7 cell media by passage over a MAb column specific for the M, 80 K portion. As shown in Table 4, approximately 65% of the applied activity was retained by the column and 50% of this bound material was eluted in an active form and at a fivefold greater concentration than in the initial media.

active complex can be isolated by affinity chromatography using an antibody specific for only the M_r 80 K species. 100 μ g of an anti-80 K monoclonal antibody (56 IgG) (Nordfang et al, Thromb Haemostasis (1985) 53:346) coupled to Sepharose® CL4B were incubated overnight at 20°C with 1.4 mL of medium containing a total of 6.2 mU of activity (measured by the COATEST Assay obtained from COS7 cells cotransfected with psvF8-92 and psvF8-80 plasmids). After incubation, the slurry was loaded into a column and the flowthrough fraction was collected. The column was washed with 300 μ L of Buffer A (50 mM imidazole, 0.1 M NaCl, 0.1% sodium insulin, 0.2% NaN₃, pH 7.3) and then eluted with 300 μ L of Buffer B (2.5 M NaCl, 50% ethylene glycol, 0.5 M imidazole, 0.1 M CaCl₂, 0.1% sodium insulin, 0.2% NaN₃, pH 7.3).

TABLE 4: Partial Purification of M, 92 K-80 K Coagulation Active Complex

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	Fraction	COATEST U/mL	80 K ELISA U/mL	
5	Media	.0044	0.175	
	Flowthrough	.0017	0.13	
	Eluate	.0200	0.76	

Results reported here demonstrate that expression of the linker ("B") region, containing 918 amino acids or about 40% of the total for the intact protein, is not required for Factor VIII:C activity. Co-expression of individual M_r 92 K and M_r 80 K regions results in a level of Factor VIII:C activity comparable to that obtained from the expression of the whole Factor VIII:C coding region. These proteins assemble in vivo to form an active complex linked by a calcium bridge. The assembly does not require the presence of the B region and occurs efficiently for the two chains expressed in trans.

It is evident from the above results that Factor VIII:C activity can be achieved by directly producing an N-terminal fragment and a C-terminal fragment which are independently expressed, each having its own signal sequence. Thus, Factor VIII:C can be obtained more efficiently, since the large precursor need not be cloned and used as the coding sequence for the Factor VIII:C activity. Thus, cells may be employed for expression of Factor VIII:C which may be deficient in the capability for proper maturation of the full-length Factor VIII:C protein.

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Example 2

Expression of the M_r 92 K protein in COS7 cells using the pSVF8-92 construction was low compared to the amount of M_r 80 K protein produced. The M_r 92 K protein is apparently retained and/or degraded in the Golgi pathway, and is not efficiently processed or exported. Accordingly, the construction was modified in an attempt to increase the level of M_r 92 K protein. Modifications of the following types were made: Changes in the 5' untranslated sequence of the Factor VIII:C gene; inclusion of heterologous 5' untranslated and leader sequences; and changes in the 3' untranslated sequences. These constructs are summarized below.

Plasmid pSVF8-92B. This plasmid is a derivative of pSVF8-92 in which the 30 bp of 5' untranslated sequence of pSVF8-92 is replaced with the entire 5' untranslated region of human Factor VIII:C cDNA (nucleotides 1 to 171; see Fig. 8 of Truett et al, supra), with a deletion of the G-C tails (by in vitro sitespecific mutagenesis), and the three base changes shown below at the starting ATG (at position +172, Fig. 8, Truett et al, supra) to conform to Kozak's preferred sequences for efficient message translation in eukaryotic cells:

Factor VIII:C: GTC<u>ATG</u> CAA
Kozak consensus: ACC<u>ATG</u> G

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This change alters the second amino acid of the signal peptide to Glu from Gln.

Plasmid pSVF8-92E: This plasmid is a derivative of pSVF8-92B in which the polylinker derived from pSV7d 5' to the Factor VIII:C sequences is removed with the exception of the SalI site, and the ATG codon in the 5' untranslated region (at 41 according to Truett et al, supra) is altered to ATT, by in vitro mutagenesis.

(B) Addition of Heterologous 5' Sequences Plasmids pSVF8-92G, H, and I. These plasmids are derivatives 15 of pSVF8-92B in which the 5' untranslated region as well as the natural Factor VIII:C signal sequences are replaced with the analogous region from the human tissue plasminogen activator (tPA) cDNA. In pSVF8-92G the first 35 amino acids (signal and pro-sequences) of the tPA pre-pro region are joined to mature 20 Factor VIII:C M_r 92 K with a serine substituted for the first amino acid (alanine) of the Mr 92 K protein. In pSVF8-92H the first 32 amino acids of the tPA pre-pro region are joined to mature Factor VIII: C M, 92 K protein. In pSVF8-92I, the first 25 23 amino acids of the tPA pre-pro region are joined to mature Factor VIII: C M, 92 K protein. The tPA sequences are the same as those described for pSVF8-80.

Plasmid pSVF8-92J. This plasmid is a derivative of pSVF8-92G in which the tPA 5' region is replaced with 75 bp of Herpes simplex virus-1 (HSV-1) gD 5' untranslated sequences and 75 bp of HSV-1 gD signal sequence. pSVF8-92J also lacks the Ala → Ser substitution (R.J. Watson et al, Science (1982) 218:381-384).

(C) <u>3' Untranslated Region Changes</u>

<u>Plasmid pSVF8-92C</u>. This plasmid is a variation of pSVF8-92B in which the M_r 92 K coding region is fused directly to the translational stop codon and natural 3' untranslated sequences of

<u>Plasmid pSVF8-92L</u>. This plasmid is a derivative of pSVF8-92C in which the 3' untranslated region of pSVF8-92C is replaced with the 3' untranslated region of pSVF8-80.

(D) <u>Results</u>

human Factor VIII:C cDNA.

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Each of the plasmids of parts A-C above was transfected into COS7 cells along with pSVF8-80 as described in Example 1 and the media tested for Factor VIII:C activity as in Example 1(F).

Plasmid pSVF8-92B, the first tested, showed activity levels ranging from 2-to-8-fold better than pSVF8-92. Of the remaining plasmids pSVF8-92E appeared to be the best, being 1.65-fold better than pSVF8-92B. pSVF8-92J and I also produced substantially higher expression levels than pSVF8-92, being close to that of pSVF8-92E. The expression level of pSVF8-92G approximated that of pSVF8-92, whereas that of pSVF8-92H was substantially less than pSVF8-92. The expression levels of both pSVF8-92C and pSVF8-92L appear to be equivalent to that of pSVF8-92E.

Example 3

This example describes the preparation of constructs for producing polypeptides that consist of the $M_{\rm r}$ 92 K chain and a portion of the B domain. These derivatives were made in an attempt to develop a heavy chain that is more stable and/or assembles more efficiently into an active complex with the light chain. The derivatives were chosen to mimic molecular species that have been observed in plasma-derived preparations

of Factor VIII:C and in cell lysates and conditioned media from cells expressing recombinant full-length Factor VIII:C. Polypeptides of approximately the same size could possibly arise by thrombin cleavages of full-length Factor VIII:C.

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- (A) pSVF8-92S: This plasmid encodes a 982 amino acid heavy chain and was prepared from a full-length cDNA plasmid pSVF8-302 by cleavage at the first SacI site of the B-domain coding region. An oligonucleotide adaptor was used to install a translational stop codon and fuse the coding sequence to the natural human Factor VIII:C 3' untranslated sequence beginning at the first Ball site. This plasmid encodes the first 978 amino acids of native human Factor VIII:C and 4 substituted amino acid residues at the carboxy terminus.
- amino acid heavy chain and was prepared from a full-length clone (designated pSVF8-303) similar to pSVF8-200, but having the 5' untranslated region of pSVF8-92E. pSVF8-303 was cleaved with EcoRV and SmaI, and the blunt ends were ligated together to form pSVF8-160. This plasmid encodes the first 1315 amino acids of Factor VIII:C. Eight substituted amino acids are added at the carboxyl terminus as a result of the fusion of the polylinker of the vector pSV7d.
- (C) pSVF8-170: This plasmid provides a 1416 amino acid heavy chain and was also prepared from pSVF8-303. pSVF8-303 was partially digested with BglII, and the resulting 6811 bp fragment was gel isolated and the ends ligated together to form pSVF8-170. This plasmid encodes the first 1405 amino acids of Factor VIII:C and has a carboxyl extension of 11 amino acids due to fusion of the polylinker of the vector pSV7d.
- (D) pSVF8-120: This plasmid provides a 1107 amino acid heavy chain and was prepared from pSVF8-303. The plasmid pSVF8-303 was digested with ApaI and the cohesive ends were filled in with T4 polymerase. The resulting molecule was further digested with SmaI, the DNA self-ligated and propagated

in *E. coli* HB101. This plasmid encodes 1102 amino acids from the amino terminus of Factor VIII:C plus an additional 5 amino acids at the carboxyl terminus, encoded by the pSV7d polylinker.

(E) Results

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Each of the plasmids of parts A-D was transfected into COS7 cells along with pSVF8-80, and the media tested for Factor VIII:C activity, as described in Example 1.

All of these plasmids showed substantially reduced expression levels compared to that of pSVF8-92E. Interestingly, though, the ratio of RIA to COATEST activity for pSVF8-160 and pSVF8-170 is about 1.8, compared to 7.2 for pSVF8-92E. This result suggests that these longer heavy chain derivatives have a higher specific activity, that is, they are more efficiently assembled into active subunit complexes than the M_r 92 K molecule itself. Also, the ratio of coagulation activity to COATEST activity is lower for the longer heavy chains at about 1.7 compared to 2.3 for M_r 92 K and 1.35 for the complete molecule, suggesting that these longer polypeptides form complexes which are not as activated as that of the M_r 92 K + M_r 80 K complex.

Example 4

This example describes the preparation of stable CHO cell lines that produce the Factor VIII:C $M_{\rm r}$ 92 K-80 K chain complex.

(A) Preparation of a plasmid encoding a selectable marker
The plasmid pAd-DHFR, bearing the murine DHFR cDNA,
was constructed by fusing the major late promoter from adenovirus-2 (Ad-MLP, map units 16-27.3) to the 5' untranslated
sequences of the mouse DHFR cDNA (J.H. Nunberg et al, Cell
(1980) 19:355-64). SV40 DNA encoding part of the early transcription unit, including the intron of the small t antigen
gene, and having the SV40 early region transcriptional term-

ination region, was obtained from pSV2-neo (Southern and Berg, <u>J Mol Appl Gen</u> (1982) 1:327-41) and fused to the 3' untranslated end of the DHFR cDNA. These three segments were subcloned into pBR322 to obtain plasmid pAd-DHFR.

(B) Transfection and culture of CHO cells

CHO-DUKX-B11 cells carrying non-functional genes for dihydrofolate reductase (Urlaub and Chasin, Proc Nat Acad Sci USA (1980) 77:4216-4220) were transfected with a calcium phosphate coprecipitate of three plasmids: pSVF8-92C, pSVF8-92E, or pSVF8-80, and pAd-DHFR following the method of Graham and Van der Eb, supra, and modifications described by Wigler et al, Cell (1978) 14:725-731 and Lewis et al, Somatic Cell Genet (1980) 6:333-347. Coprecipitates contained up to 10 μ g of each plasmid. Cells were selected for expression of the DHFR (positive) phenotype in a medium deficient in hypoxanthine and thymidine.

After isolation of DHFR positive clones and identification of those producing Factor VIII:C activity, the resulting cell lines were grown in methotrexate to amplify the DHFR genes and coamplify the Factor VIII:C genes. This selection was performed by plating cells in medium containing methotrexate in concentrations ranging from 0.025 to 0.2 μ M. Methotrexate resistant clones were again assayed for Factor VIII:C activity.

(C) Assay Methods

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Conditioned media from these DHFR positive clones were assayed by ELISA for Factor VIII:C light chain immunoreactivity by the method of Nordfang et al, Thromb Haemostas (1985) 53:346-50. Factor VIII:C heavy chain immunoreactivity was evaluated using a radioimmunoassay (RIA) described by R.L. Burke et al, J Biol Chem (1986) 261:12574-78. Active Factor VIII:c complexes formed by co-expression of the 92 K and 80 K M, glycoproteins were measured using the COATEST assay described in Example 1.

(D) CHO lines expressing active 92 K-80 K Mr complexes Shown in Table 5 are four independent CHO cell lines that simultaneously express products of all three plasmids used for transfection. The Factor VIII:C activity values shown in Table 5 are those initially observed. Expression of glycoproteins by stable cell lines usually improves after passage in T-75 flask cultures. An example of this can be seen for the line 10-C2, which ultimately produced 200 mU Factor VIII:C activity per mL conditioned medium (Table 6). Cloning these stable cell lines illustrates that the independently expressed heavy and light chains of Factor VIII:C can assemble into an active complex and be secreted by Chinese hamster ovary cells.

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CHO cell lines producing active 92 K-80 K complexes TABLE 5:

Clone	one Transfected DNA			mU COATEST/mL
11-D6 11-D5	pSVF8-92C, pSVF8-92C,	pSVF8-80,	pAd-DHFR	· 43 30
8-C1 10-C2	pSVF8-92E, pSVF8-92E,			18.2 70.0

. That the three plasmids were integrated into the chromosomes of the CHO cells is suggested by the fact that the cell lines of Table 5 could be grown for many passages without loss of Factor VIII:C expression. It was then necessary to determine if expression of Factor VIII:C glycoproteins could be co-amplified by methotrexate selection. All four of these cell lines were placed under selection in several concentrations of methotrexate. Resistant colonies (DHFR genes amplified) were obtained for each line and these were screened for Factor VIII:C activity. Expression of Factor VIII:C was lost or unchanged in methotrexate resistant 11-D5 and 11-D6 clones. Expression of Factor VIII:C varied among methotrexate resistant clones derived from 10-C2 and 8-C1 (shown in Table 6).

Twenty-two methotrexate-resistant 8-C1 clones were examined, the data for 10 of which are reported in Table 6.

The amount of Factor VIII:C amplification varies among clones, suggesting that either one of the subunit genes may have been co-amplified with the DHFR cassette, or both of them, or neither one. Note clones 8C1-A2, 8C1-C2, and 8C1-C5 as examples of these four possibilities. Similarly, 30 methotrexate-selected derivatives of 10-C2 were evaluated, the data for 20 of which are also represented in Table 6. These also contain a spectrum of activity. Note clones 10C2-A2, 10C2-D2, 10C2-B5, and 10C2-C6 as examples of the four different co-amplification possibilities.

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TABLE 6

5	Clone	conc. MTX (µM)	COATEST (mU/mL)	LC-ELISA (mU/mL)	HC-RIA (mU/mL)
5	8-C1	0	18	1275	n.d.
	8C1-A1	0.1	<50	1750	80
	8C1-A2	0.1	60	1950	>1000
	8C1-A5	0.05	. 2	100	10
10	8C1-B3	0.025	33	1950	1000
	8C1-B4	0.025	50	3550	820
	8C1-B5	0.025	35	1950	>1000
	8C1-C2	0.025	130	13,100	>>1000
	8C1-C3	0.025	165	3900	>>>1000
15	8C1-C5	0.025	30	1750	760
	10-C2	0	200	1400	700
	10C2-A1	0.05	61	1600	. 400
	10C2-A2	0.1	67	6700	700
	10C2-A4	0.05	63	2250	1200
20	10C2-A5	0.05	183	9450	2660
	10C2-A6	0.05	320	8600	7400
	10C2-B1	0.05	408	8100	4300
	10C2-B3	0.05	134	800	9800
	10C2-B4	0.05	394	18,000	7800
25	10C2-B5	0.05	461	15,000	8400
	10C2-B6	0.05	247	2200	9800
	10C2-C1	0.1	160	8100	7600
	10C2-C2	0.05	228	6000	5600
	10C2-C3	0.05	294	14,850	2650
30	10C2-C5	0.05	294	12,400	5400
	10C2-C6	0.05	100	1350	520
	10C2-D2	0.05	496	1560	16,400
	10C2-D3	0.05	242	10,200	2260
	10C2-D4	0.05	165	14,100	3500
35	10C2-D5	0.05	316	7800	5200
	10C2-D6	0.05	141	1600	6400

Among the CHO lines described in Table 6 is one

(10C2-D2) that produces 0.5 U/mL of active Factor VIII:C complex, which is one half the concentration found in normal human plasma. For analysis and purification of Factor VIII:C material, CHO cell lines expressing Factor VIII:C polypeptides were grown in laboratory scale fermentations to produce 1-2 liter quantities of tissue culture fluid. Assay of this material showed that approximately 10% to 20% of immunoreactive Factor

VIII:C from unamplified lines is active in the COATEST. In amplified lines, the percentage of active material drops to 2% to 5% of the total immunoreactive product. This means that only a fraction of the heavy and light chains of FVIII:C is assembled into active complexes. The remainder may exist as free subunits or in degraded forms.

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Plasmids pSVF8-92 and pSVF8-80 were deposited at the American Type Culture Collection (ATCC) on 24 January 1986 and given ATCC Accession Nos. 40222 and 40223, respectively. Plasmid pSVF8-200 was deposited at the ATCC on 17 July 1985 and was given ATCC Accession No. 40190.

Example 5

This example describes modification of the plasmid pSVF8-80 to correct the amino terminal amino acid of the FVIII:C light chain glycoprotein. A consequence of engineering, which provided the signal peptide needed for independent secretion of the 80 K M_r glycoprotein (Example 1) is the substitution of Ser for the normal aminoterminal residue of human plasma FVIII:C light chains. New plasmids were made in an attempt to change the tPA pre-pro peptide sequence, so that the FVIII:C light chain will have the Glu residue at its amino terminus instead of the mutant Ser residue after proteolytic processing.

The FVIII:C light chain is thought to be cleaved from the full-length FVIII:C precursor before secretion, i.e. intracellularly, by a protease resident in the Golgi apparatus. This cleavage occurs between amino acid residues 1648 and 1649 (Arg-Glu). On polyacrylamide gels the light chains appear as a doublet of 77 and 80 K M_r bands, representing polypeptides having one or two N-linked oligosaccharides. Independent secretion of light chains was achieved by fusion of the light chain coding region of the FVIII:C cDNA to the cDNA of tPA. In the process of supplying the tPA signal peptide, however, the amino

terminus of the FVIII:C light chain was mutated from the native glutamic acid residue to a serine. Although this mutant recombinant light chain displays molecular characteristics similar to the chain derived from full-length recombinant FVIII:C, there is preliminary evidence that 1) it may not be alternatively glycosylated in the same manner as the chain cleaved from the FVIII:C precursor, 2) it may behave differently during purification by ion exchange and vWF Sepharose® chromatography, and 3) it may be different antigenically from authentic light chain.

The tPA pre-pro peptide sequence requires three protectly cleavages to release the mature polypeptide. Shown below is the translation of the protein coding sequence of pSVF8-80 in the region of the tPA-FVIII:C 80 K fusion:

pSVF8-80:

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-35 -30 -25

Met Asp Ala Met Lys Arg Gly Leu Cys Cys Val Leu Leu Leu

20 ATG GAT GCA ATG AAG AGA GGG CTC TGC TGC TGT GTG CTG

-20 -15 * -10 *

Cys Gly Ala Val Phe Val Ser Pro Ser Gln Glu Ile His Ala

TGT GGA GCA GTC TTC GTT TCG CCC AGC CAG GAA ATC CAT GCC

-5 0 1 5
Arg Phe Arg Arg Gly Ala Arg Ser Ile Thr Arg Thr Thr Leu
CGA TTC AGA AGA GGA GCC AGA TCT ATA ACT CGT ACT CTT CAG

Gln Ser Asp 35 CAG TCT GAT

The signal peptidase cleavage has been thought to occur on the carboxy side of either Ser (position -13) or Ala (position -8), indicated by asterisks. The second cleavage probably occurs on

the carboxy side of Arg (position -4, indicated by @ above). The third processing event is proteolysis at the Arg-Ser bond to release a Gly-Ala-Arg tripeptide and leave a Ser (position 1) aminoterminus on the mature tPA or FVIII:C light chain polypeptides.

(A) Preparation of plasmids

(1) pSVF9-80KG:

ted mutagenesis to a Glu codon (position 1). This was done in an effort to allow the first two proteolytic processing events to occur normally, and test whether the Arg-Glu protease could recognize and cleave the dipeptide in an altered context, i.e., where the tPA tripeptide is substituted for the FVIII:C B domain. The tPA-80 K chain fusion region is shown below.

Otherwise, this plasmid is identical to pSVF8-80.

pSVF8-80KG:

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20 Met Asp Ala Met Lys Arg Gly Leu Cys Cys Val Leu Leu ATG GAT GCA ATG AAG AGA GGG CTC TGC TGC TGT GTG CTG 25 -20 -15 -10 Cys Gly Ala Val Phe Val Ser Pro Ser Gln Glu Ile His Ala TGT GGA GCA GTC TTC GTT TCG CCC AGC CAG GAA ATC CAT GCC 30 Arg Phe Arg Arg Gly Ala Arg Glu Ile Thr Arg Thr Thr Leu CGA TTC AGA AGA GGA GCC AGA GAA ATA ACT CGT ACT CTT CAG 35 10 Gln Ser Asp CAG TCT GAT

(2) pSVF8-80S:

Twelve codons were deleted from pSVF8-80 by in vitro mutagenesis, and the Ser (position 1) codon changed to a codon for Glu. This placed the Glu FVIII:C light chain residue after Ser23 of the putative tPA signal peptide (indicated by an asterisk). Cleavage by signal peptidase on the carboxy side of Ser23 releases the non-mutant FVIII:C light chain. The tPA - 80 K chain fusion region of pSVF8-80S is shown below. Otherwise this plasmid is identical to pSVF8-80.

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pSVF8-80S:

-23 -20 -15 -10

Met Asp Ala Met Lys Arg Gly Leu Cys Cys Val Leu Leu Leu

ATG GAT GCA ATG AAG AGA GGG CTC TGC TGC TGT GTG CTG CTG

-5 * 1 5

Cys Gly Ala Val Phe Val Ser Pro Ser Glu Ile Thr Arg Thr

TGT GGA GCA GTC TTC GTT TCG CCC AGC GAG ATA ACT CGT ACT

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Thr Leu Gln Ser Asp Gln Glu Glu Ile Asp Tyr Asp Asp Thr CTT CAG CAG TCT GAT CAA GAG GAA ATT GAC TAT GAT GAT ACC

(3) pSVF8-80R:

A deletion of three codons of pSVF8-80, to remove the tPA pro-tripeptide, was made by in vitro mutagenesis, and the Ser (position 1) codon was changed to one for Glu. This places a Glu residue after Arg32 of the tPA pro-peptide, marked with @ on the tPA-80 K chain fusion region of pSVF8-80R shown below:

pSVF8-80R:

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-32 -30 -25 -20

Met Asp Ala Met Lys Arg Gly Leu Cys Cys Val Leu Leu Leu

5 ATG GAT GCA ATG AAG AGA GGG CTC TGC TGC TGT GTG CTG

-15 -10 -5
Cys Gly Ala Val Phe Val Ser Pro Ser Gln Glu Ile His Ala
10 TGT GGA GCA GTC TTC GTT TCG CCC AGC CAG GAA ATC CAT GCC

0 1
Arg Phe Arg Arg Glu Ile Thr Arg Thr Thr Leu Gln Ser Asp
CGA TTC AGA AGA GAG ATA ACT CGT ACT CTT CAG CAG TCT GAT

This construction was made in the hope that cleavage by a Golgi-resident protease with dibasic specificity would release FVIII:C light chains having Glu amino termini.

(4) pSVF8-80A:

Seven codons of pSVF8-80 were deleted by site-directed mutagenesis, removing the DNA encoding the putative tPA pro sequence, and the Ser (position 1) codon was replaced by a Glu codon after codon 28 (Ala) of the putative tPA signal peptide coding sequence (indicated by an asterisk below). Cleavage by signal peptidase on the carboxy side of Ala₂₈ will release non-mutant FVIII:C light chain. The tPA-80 K chain fusion region is shown below. Otherwise, this plasmid is identical to pSVF8-80.

pSVF8-80A:

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-28 -25 -20 -15
Met Asp Ala Met Lys Arg Gly Leu Cys Cys Val Leu Leu Leu
ATG GAT GCA ATG AAG AGA GGG CTC TGC TGC TGT GTG CTG

-10 -5 *
Cys Gly Ala Val Phe Val Ser Pro Ser Gln Glu Ile His Ala
TGT GGA GCA GTC TTC GTT TCG CCC AGC CAG GAA ATC CAT GCC

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Glu Ile Thr Arg Thr Thr Leu Gln Ser Asp Gln Glu Glu Ile
GAG ATA ACT CGT ACT CTT CAG CAG TCT GAT CAA GAG GAA ATT

(B) Expression and protein sequence analysis

(1) Transfection into COS7 Cells:

COS7 cells were transfected using the DEAE-dextran procedure described in Example 1, and conditioned media were assayed by the LC-ELISA. All four derivatives of pSVF8-80 encode 80 K M_r glycoproteins that are reactive in the LC-ELISA and that can be immunoprecipitated after biosynthetic radio-labeling with various anti-FVIII:C light chain antibodies. Except for pSVF8-80R, all the derivatives lead to secretion of about the same amount of 80 K glycoprotein as pSVF8-80. Secretion of 80 K glycoprotein from cells transfected with pSVF8-80R is very poor, usually less than 25% of that produced from the other plasmids. In addition, the appearance of this FVIII:C light chain is different on gel electrophoresis, where the bands are always diffuse.

(2) <u>Expression in CHO Cells</u>:

Each of these plasmids was introduced into DUKX-B11 CHO cells with pAd-DHFR as described in Example 4. Permanent cell lines were established for production of each type of light chain. Expression of the 80 K M_r glycoproteins in CHO cells is very similar to expression in COS7 cells, with respect

to the amounts of glycoprotein secreted and the appearance of the 80 K bands on gel electrophoresis. CHO lines transfected with pSVF8-80R produced such a low level of 80 K glycoprotein that analysis of this material was not done.

3. <u>Purification and amino acid</u>

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Conditioned media from either large scale COS7 transfections (pSVF8-80KG) or from transfected (amplified) CHO cell lines (pSVF8-80K cell line 10C2B5; pSVF8-80A, cell line A1N; pSVF8-80S, cell line S1R) were prepared. The medium was DME H12 with 10% FBS. FVIII-LC was purified for sequencing by a two-step procedure comprising ion exchange chromatography followed by affinity chromatography. Ion exchange chromatography was performed as follows: A column of S-FF Sepharose® (15 × 0.8 cm) was equilibrated with 0.02 M MES, 0.05 M NaCl, 0.01 M CaCl₂, pH 5.8, $\lambda_{20^{\circ}C} = 7.2$ mS. Conditioned medium (500-1300 mL) was applied to the column after adjustment of pH to 5.8 with a flow rate of 100 mL/h. The column was washed with 10 column volumes of 0.05 M imidazole, 0.05 M NaCl, 0.01 M CaCl, pH 7.35, $\lambda_{20^{\circ}C} = 8.8$ mS at a flow rate of 200 mL/h. FVIII-LC was eluted by addition of 0.1 M CaCl, to the washing buffer, flow rate 50 mL/h. All operations were performed at 4°C.

Affinity chromatography was performed as follows: The murine monoclonal anti-FVIII-LC antibody 56-IgG was coupled to Sepharose® 4B by the CNBr method to a density of 2.5 mg/mL gel. The FVIII-LC containing eluate was incubated with the immunosorbent overnight at room temperature, 1 mL of gel per 1000 units FVIII-LC. The gel was then packed into a column and washed with 20 column volumes of a low salt buffer (0.05 M imidazole, 0.15 M NaCl, 0.01 M CaCl₂, 10% glycerol, 0.02% NaN₃, pH 7.3), followed by 20 column volumes of a high salt buffer (0.05 M imidazole, 1.0 M NaCl, 10% glycerol, pH 7.3). FVIII-LC was eluted from the immunosorbent using 1 M CaCl₂ in 0.05 M

imidazole, 0.15 M NaCl, 10% glycerol after one hour incubation. The eluate was immediately desalted on a Sephadex® G-25 column to a solution of 0.05 M imidazole, 0.15 M NaCl, 0.01 M CaCl₂, 10% glycerol, 0.02% Tween® 80, 0.02% NaN₃, pH 7.3 and stored at -80°C. N-terminal sequence analysis was performed on an Applied Biosystem 477A sequencer.

The results of this analysis are shown in Table 7. The 80 K glycoprotein encoded by pSVF8-80KG has a tripeptide extension on its aminoterminus. Presumably this is the tPA pro tripeptide Gly-Ala-Arg, which cannot be processed by the Arg-Glu protease that recognizes the FVIII:C B domain. Further, the N-terminal sequences reveal that the signal peptide of tPA is actually 22 amino acid residues in length, with signal peptidase cleavage occurring on the carboxy side of Pro₂₂. Therefore, plasmid constructions pSVF8-80S and pSVF8-80A, predicated upon signal peptidase cleavage after Ser₂₃ and Ala₂₈, respectively, lead to incorrect amino terminal residues on the 80 K light chains.

TABLE 7: N-terminal Sequences of 80 K Chains with Modified tPA pre-pro Regions

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Plasmid	N-terminal Sequence	Amount (pmol)
pSVF8-80	X-Ile-X-Arg-Thr-X-Leu-Gln-X-Asp-Gln-	10
pSVF8-80KG	X-X-Arg-Glu-Ile-Thr-Arg-Thr-Thr-Leu-	20
pSVF8-80S	Ser-Glu-Ile-Thr-Arg-Thr-	40
pSVF8-80A	X-Gln-Glu-Ile-	. 40

Results shown in this example reveal the difficulty of predicting how a secreted polypeptide will be processed following transcription and translation. Modifications of the protein sequence have unexpected consequences for proteolytic

processing and oligosaccharide addition, and can affect the overall efficiency of secretion.

Example 6

This example describes a method for expression of authentic FVIII:C light chains using the signal peptide of human α_1 -antitrypsin.

A. Preparation of plasmids

pSVα1AT.Met

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A cDNA encoding the mature human α_1 -antitrypsin polypeptide had been assembled using fragments of human liver cDNA clones and a synthetic oligonucleotide; the assembly was ligated as a BamH1-SalI fragment into pBR322 to make plasmid pAT(Met) (Rosenberg et al, Nature (1984) 312:77-80). A synthetic oligonucleotide linker-adapter and part of a cDNA clone encoding the signal peptide were used to attach the signal peptide coding sequence, with an EcoRI restriction site on the 5' end, to the BamHI site of pAT(Met). The resulting 1271 bp EcoRI-SalI fragment, encoding the translated sequences of human α_1 -antitrypsin, was ligated into the EcoRI-SalI sites of pSV7d (described in Example 1) to make pSV α 1AT.Met.

2. pSVF8-80AT

Plasmid pSV α lAT.Met was opened at the BamHI site, which occurs at the boundary between the codons of the signal peptide and mature α_1 -antitrypsin sequences. The cohesive end of this restriction site was removed with mung bean nuclease to leave the GAG (Glu) codon, and the α_1 -antitrypsin sequence was deleted by digestion with SalI. The coding sequence of FVIII:C 80 K was prepared for attachment by in vitro mutagenesis of codons 1 and 2 of pSVF8-80 to form an EcoRV site (which preserves codon 2 as an Ile codon). This allowed the FVIII:C light chain coding sequence (as an EcoRV-SalI sequence starting at codon 2) to be fused in correct reading frame to codon 1 of

 α_1 -antitrypsin, and replace the coding sequence of mature human α_1 -antitrypsin.

The coding sequence of pSVF8-80AT in the region of fusion is shown translated below. Except for substitution of the α_1 -antitrypsin signal peptide coding sequence for the tPA pre-pro coding sequence, this plasmid is identical to pSVF8-80.

والأركاز ويرزي والمتأثث والمتأثث والمناكل والموجول والمناك المسروي بالتعاجب والروائد والمثان والمتحر والمتحر والمتاكا

pSVF8-80AT (amino terminal region):

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Met Pro Ser Ser Val Ser Trp Gly Ile Leu Leu Ala Gly Leu

ATG CCC TCG AGC GTC TCG TGG GGC ATC CTC CTG CTG GCA GGC CTG

Cys Cys Leu Val Pro Val Ser Leu Ala Glu Ile Thr Arg Thr Thr TGC TGC CTG GTC CCT GTC CTG GCT GAG ATC ACT CGT ACT

Leu Gln Ser Asp Gln Glu Glu Ile Asp Tyr Asp Asp Thr Ile Ser
CTT CAG TCT GAT CAA GAG GAA ATT GAC TAT GAT ACC ATA TCA

B. <u>Expression and amino acid sequence analysis</u>

1. Expression of pSVF8-80AT in COS7 cells

heavy chain expression plasmid, usually pSVF8-92C. Conditioned media were assayed by LC-ELISA, HC-ELISA and COATEST. Transfected cells were also labeled with radioactive Met, so that the biosynthetically radiolabeled FVIII:C light chains could be immunoprecipitated and visualized after polyacrylamide gel electrophoresis. Plasmid SVF8-80AT directs the synthesis of FVIII:C light chains that appear as a doublet of 77-80 K M_r. The amount produced in COS7 cells is the same as for pSVF8-80. Co-expression with pSVF8-92C, or other FVIII:C heavy chain plasmid, leads to production of active FVIII:C complexes measured in the COATEST assay.

2. Purification and amino acid sequence analysis

Material for purification was prepared by transfection of COS7 cells in T-175 flasks, using increased cell density and decreased chloroquine diphosphate concentration. Conditioned media were collected 60 hours after transfection.

Purification and amino acid sequence analysis were performed as described in Example 5. The results of aminoterminal sequence analysis (Table 8) indicate that the FVIII:C light chain encoded by pSVF8-80AT has the same aminoterminal sequence as authentic human plasma FVIII:C light chain.

TABLE 8: N-terminal Sequence of 80 K Chains Secreted Using α_1 -antitrypsin Signal Peptide

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15 .	Plasmid	N-terminal Sequence	Amount (pmol)
	pSVF8-80	X-Ile-X-Arg-Thr-X-Leu-Gln-X-Asp-Gln-	10
20	pSVF8-80AT	Glu-Ile-Thr-Arg-Thr-X-Leu-Gln-Ser-Asp-Gl	n 10

3. In vitro assembly of 80AT FVIII: C light chains

The ability of 80 AT FVIII:C light chains to recombine with purified FVIII:C heavy chains in vitro was tested in an experiment shown in Table 9. Purified FVIII:C light chains were incubated at concentrations of 3.7 U/mL with purified recombinant (from full-length human FVIII:C) heavy chains at 17 U/mL in buffer containing 50 mM Mn $^{++}$ and 150 μ M β -mercaptoethanol. As the control, purified recombinant heavy and light chains were allowed to reassociate under the same conditions, and the quantity of active FVIII:C produced was assayed by COATEST. These results suggest that the 80AT FVIII:C light chain can be combined in vitro with purified recombinant heavy chain.

TABLE 9: Combination of Recombinant FVIII:C Light Chains with Heavy Chains in vitro

Purified FVIII-LC	Purified FVIII-ḤÇ	Percent Control Activity	
TA08	full-length	86	
805	full-length	51	
80A	full-length	. 92 .	
80KG	full-length :	92 	··· ·

Example 7

This example describes plasmids for improved expression of the Factor VIII:C heavy chain. Modifications in DNA sequences responsible for the initiation of transcription and in non-coding sequences are made in order to increase the efficiency of transcription and the stability of the messenger RNA. The heavy chain glycoprotein is modified by a carboxy-terminal extension composed of segments of the B domain joined by a short peptide. This is done to obtain a heavy chain that is secreted from cells more efficiently, is more stable in tissue culture medium, and assembles more efficiently with the light chain.

A. Preparation of plasmids

1. pCMVF8-92/6x

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In an effort to improve the level of transcription and stability of the messenger RNA for the Factor VIII:C 92 K M_r heavy chain, the SV40 early transcriptional initiation region was replaced by sequences from the human cytomegalovirus immediate early region (Boshart et al, Cell (1985) 4:521-530). In addition, 5' untranslated sequences contributed by the SV40 early region to the messenger RNA were replaced with the 5' untranslated sequences of the HCMV 1E1 gene, including its first intron. This intron is included on the assumption that spliced transcripts lead to faster processing and more stable

mRNA. The expression vector also has an SV40 origin of replication to permit transient expression in COS7 cells, and a bacterial β -lactamase gene to permit DNA cloning by selection for ampicillin resistance.

The plasmid was constructed from a 700 bp SalI-PvuI fragment of pSV7d (described in Example 1) containing the SV40 polyadenylation region, a 1400 bp PvuI-EcoRI (filled in with Klenow polymerase) fragment of pSVT2 (Myers et al, Cell (1981) 25:373-84; Rio et al, Cell (1983) 32:1227-40) providing the SV40 origin of replication and the rest of the β -lactamase gene, a 1700 bp SspI-SalI fragment derived from a plasmid subclone of the human cytomegalovirus (Towne strain) in which the SalI site was introduced by in vitro mutagenesis near the translational start site for the 1E1 protein, and the 4300 bp SalI-SalI fragment of pSVF8-92C (described in Example 2) containing the cDNA encoding the Factor VIII:C 92 K Mr glycoprotein.

2. $pSVF8-92t\beta$

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This plasmid is a derivative of pSVF8-92C that encodes the 92 K M, recombinant heavy chain with a C-terminal extension composed of N-terminal and C-terminal amino acid residues of the central (B) domain of the Factor VIII:C precursor linked by a peptide hinge peptide homologous to that of 25 human immunoglobulin α heavy chain. It is composed of a 4900 bp HindIII-SalI fragment from pSVF8-92C, into which was inserted a 110 bp HindIII-SalI synthetic linker-adapter (shown below).

N-terminus of B Domain
SerPheSerGlnAsnSerArgHisProSerThrArgGlnLysGlnPheAsnAla AGCTTCTCCCAGAATTCTAGACACCCTAGCACTAGGCAAAAGCAATTTAATGCC
AGAGGGTCTTAAGATCTGTGGGATCGTGATCCGTTTTCGTTAAATTACGG

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ThrProProThrProProThrProProValLeuLysArgHisGlnArgOP OC ACCCTCCTACACCACCACCACCACCAGTACTGAAACGCCATCAACGG<u>TGATAA</u>G TGGGGAGGATGTGGTGGTTGGGGTGGTCATGACTTTGCGGTAGTTGCCACTATTCAGCT | Scall Sall

The linker-adapter encodes a carboxy-terminal extension of 34 additional amino acid residues, and one potential site of N-linked glycosylation. The C-terminal peptide should increase the molecular weight of the heavy chain to approximately 96 K Mr, and to about 99 K Mr if it is glycosylated.

25 B. Assay for FVIII:C heavy chain antigen and FVIII:C complex formation

The cofactor activity of the FVIII:C light chain-heavy chain complex was estimated using a commercially available test kit from KabiVitrum (COATEST). Immunoreactive FVIII:C light chain was measured by ELISA using HZ IgG coating antibody and peroxidase-conjugated antibodies from Nordisk Gentofte. The FVIII:C heavy chain immunoreactivity was quantified using an ELISA developed at Nordisk Gentofte, which employs human polyclonal antibody from an inhibitor patient (E-IgG).

C. Transient expression of pCMVF8-92/6x

The pCMVF8-92/6x plasmid was cotransfected with various FVIII:C light chain plasmids (described in Example 5) into COS7 cells using the DEAE-dextran procedure. A sample of

results from these transfections is shown in Table 10. The data suggest that addition of the CMV 1E1 promoter/enhancer, and the 5' untranslated sequences of the 1E1 gene yields a 2.5 fold improvement (on average) in FVIII:C heavy chain expression.

TABLE 10: Expression in COS7 Cells of pCMVF8-92/6x Versus pSVF8-92C

		C Activity	(mU/mL)
HC-Plasmid	LC-Plasmid	COA ^a	HC-RIAD
pSVF8-92C	pSVF8-80	46	160
pSVF8-92C	-80A	34	72
pSVF8-92C	-80R	61	310
pSVF8-92C	- 80S	31	46
pCMVF8-92/6x	- 80 .	87	290
pCMVF8-92/6x	-80A	131	140
pCMVF8-92/6x	-80R	178	330
pCMVF8-92/6x	-80S	114	690

COATEST assay

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D. Transient expression in COS7 Cells of pSVF8-92tß

Shown in Table 11 are the results of cotransfecting pSVF8-92t β with a Factor VIII:C light chain expression plasmid (pSVF8-80AT, described in Example 6) into COS7 cells. The 92t β heavy chain is secreted at higher levels than the 92C heavy chain, which has a single amino acid (Ser) carboxy-terminal extension. The ratio of COATEST (COA) activity to ELISA-reactive glycoprotein (a measure of complex formation) is greater for 92t β chains than for 92C chains. In addition, the 92t β heavy chain is secreted well in serum-free medium and appears to be stable, with a ratio of activity to protein nearly the same as in 10% FBS. These results show that this 34 amino acid carboxy-terminal extension improves secretion and stabilizes the recombinant FVIII:C heavy chain.

b Radioimmunoassay for heavy chain

TABLE 11: Expression of pSVF8-92t β in COS7 Cells

Exp	† Medium	Plasmid 1	Plasmid 2	mU/mL COATEST	FVIII · LC ^a	HCb
1	10% FBS	pSVF8-92tβ pSVF8-92C	psvř8–80AT psvF8–80AT	41 23	487 442	98 49
2	10% FBS	pSVF8-92tβ pSVF8-92C	pSVF8-80AT pSVF8-80AT	80	484 639	162 90
3	нв сно	pSVF8-92tβ	pSVF8-80AT	38	262	125

COS7 cell monolayers in duplicate were exposed to DNA in DEAE-Dextran, washed, and treated with medium containing chloroquine diphosphate for 8 hrs. Cells were washed to remove the drug, then covered with 5 mL DME H21 containing 10% FBS. For expression in serum-free medium, dishes were washed after 12-16 hrs. and overlaid with HB CHO® from Hana Biologicals. Conditioned media were assayed for FVIII:C activity as des-

- by ELISA assay specific for light chain.
- by ELISA assay specific for heavy chain.

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Example 8

This example describes a method for expression of a FVIII: C heavy chain having Arg, as the C-terminus.

Preparation of plasmid pCMVF8-92R Α.

The FVIII: C heavy chain encoded by the plasmid pcMVF8-92/6x has Ser741 as a C-terminal extension. In order to obtain a FVIII: C heavy chain with Arg, as the C-terminus, a 1588 bp BamHI fragment of pCMVF8-92/6x , encoding the 3' end of the coding sequence derived from pSVF8-92C was purified. fragment was cloned into m13mp18 and the Ser741 residue was changed to a translational stop codon by in vitro mutagenesis. The pCMVF8-92R expression plasmid was assembled by cloning the mutagenized BamHI fragment into the 5840 bp BamHI fragment of the original vector. By this procedure 680 bp of FVIII:C 3' untranslated sequences were deleted.

B. Transient expression in COS7 cells of pCMVF8-92R

The pCMVF8-92R plasmid was co-transfected with the FVIII:C light chain plasmid pSVF8-80AT (described in Example 6) into COS7 cells using the calcium phosphate technique (Graham and van der Eb, <u>Virol</u> (1973) <u>52</u>:456-67). The media were changed 18 and 42 hours post-transfection. Media samples for assays were collected 66 hours post-transfection. The results from these assays are shown in Table 12 below. The data shows that FVIII:C activity was generated when pCMVF8-92R was cotransfected with a plasmid providing expression of FVIII LC.

TABLE 12: Coexpression of pCMVF8-92R and pSVF8-80AT.

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Transfection	COA (mU/mL)	HC:Ag (mU/mL)	LC:Ag (mU/mL)
A	243	400	990
В	263	460	1190

Example 9

This example describes the preparation of a stable CHO cell line that produces the native FVIII:C $M_{\rm r}$ 92K-80K complex.

The DHFR CHO cell line DG44 (G. Urlaub et al, Som Cell Mol Genet (1986) 12:555-66) was first transfected with the plasmid pCMVF8-80AT. In this plasmid the CMV promoter (described in Example 7) regulates FVIII LC cDNA derived from pSVF8-80AT (described in Example 6), and downstream contains the Ad-MLP/dhfr cassette derived from pAd-DHFR (described in Example 4). The cells were transfected using the polybrene method described by W. Chaney et al, Som Cell Mol Genet (1986) 12:237-44. By selecting for DHFR+ cells (in DMEM + 10% DFSC) several FVIII-LC producers were isolated; one of which was designated 11W.

In order to introduce FVIII-HC into 11W, the cell line was cotransfected with the plasmid pPR78 (this plasmid is analogous to pCMVF8-80AT, but contains the HC cDNA derived from pCMVF8-92R described in Example 8 instead of the LC cDNA) and pSV2-neo (P.J. Southern and P. Berg, J Mol Appl Gen (1982) 1: 327-41). The transfection method used was the modified calcium phosphate procedure (G. Chen and H. Okayama, Mol Cell Biol (1987) 7:2745-52). Transfectants were isolated in medium containing 700 μ g/mL Geneticin (G418 sulfate, Gibco). Cells from the primary pool were subcloned by limiting dilution, and individual clones tested for expression of active FVIII:C. In this way several FVIII:C producing cell lines were isolated, one of which was designated 45-4/B-9.

45-4/B-9 was grown to confluency in a T-75 culture

flask (DMEM + 10% DFCS + 700 μg/mL Geneticin) at 37°C, after
which the cells were transferred to 27°C and acclimatized for 3
days before an expression period of 24 hours. After the
expression period, the FVIII:C concentration was measured to
12.8 U/mL using the chromogenic assay (Kabi).

20 Deposit Information:

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The following materials were deposited with the American Type Culture Collection:

1111022000	• • • •	Accession No.
Plasmid	Deposit Date	ACCESSION NO.
pSVF8-92	24 January 1986	40222
pSVF8-80	24 January 1986	40223
pSVF8-200	17 July 1985	40190

The above materials have been deposited with the American Type Culture Collection, Rockville, Maryland, under the accession numbers indicated. This deposit will be maintained under the terms of the Budapest Treaty on the International Recognition of the Deposit of Microorganisms for purposes of Patent Procedure. The deposits will be maintained for a period of 30 years following issuance of this patent, or for

the enforceable life of the patent, whichever is greater. issuance of the patent, the deposits will be available to the public from the ATCC without restriction.

These deposits are provided merely as convenience to those of skill in the art, and are not an admission that a The sequence of the deposit is required under 35 U.S.C. §112. polynucleotides contained within the deposited materials, as well as the amino acid sequence of the polypeptides encoded thereby, are incorporated herein by reference and are controlling in the event of any conflict with the written description 10 of sequences herein. A license may be required to make, use, or sell the deposited materials, and no such license is granted hereby.

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Although the foregoing invention has been described in some detail by way of illustration and example for purposes 15 of clarity of understanding, it will be obvious that certain changes and modifications may be practiced within the scope of the appended claims.